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(54) **THERMOSTABLE XYLANASES**

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XYLANASES THERMOSTABLES

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Description

[0001] The present invention relates to thermostable xylanase enzymes. More specifically, the present invention is directed to thermostable xylanase enzymes that exhibit high activity at or near physiological pH and temperature, and their use in feed pelleting applications.

BACKGROUND OF THE INVENTION

[0002] Natural xylanase enzymes, such as that of the fungus *Trichoderma reesei*, have been added to animal feed to increase the efficiency of digestion and assimilation of nutrients. During digestion of feed grains such as wheat and barley, non-starch polysaccharides, including xylan, increases the viscosity of the digesta in the absence of added exogenous enzyme. This interferes with the diffusion of the digestive enzymes to the feed and the subsequent assimilation of the nutrients. The highly viscous digesta increases the occurrence of sticky stool, which increases the likelihood of disease and causes effluent run-off problems. The addition of xylanase in feed breaks down the xylan and decreases the viscosity of the digesta, thereby helping to alleviate these problems. Xylanase produces a cost saving by increasing the efficiency of feed conversion. Xylanase can decrease the feed consumed/weight gain ratio by 5-15% (Viveros, A., Brenes, A., Pizarro, M. and Castano, M., 1994. *Animal Feed Sci. Technol.* 48:237-251).

[0003] Xylanase enzymes used for feed are typically aqueous solutions of active protein, stabilizers, preservatives and other additives. The enzymes are typically sprayed onto the feed at concentration of 100-2000 ml per tonne feed. Alternatively, granular or powdered xylanase can be used. Once the feed is consumed by the animal, the enzyme acts on xylan as the feed is ingested and digested in the gut. Eventually the xylanase, a protein molecule, is hydrolysed by the digestive enzymes (proteases) into amino acids like any protein in the feed.

[0004] Increasingly, animal feeds are pelleted at high temperatures for sterilization against harmful bacteria, for example *Salmonella*. Feed pelleting is carried out by heating the feed solids with 100 to 140°C steam and passing them through an extruder/pelleting auger to form feed pellets, which then cool in a storage bin. The typical time required for the material to pass through the system is 30 minutes. As is known in the art, higher temperatures can be used with shorter pelleting times, and lower temperatures with longer pelleting times, provided that the necessary moisture levels are obtained. The overall resulting temperature within the solids, prior to, during, and after pellet formation reaches about 70-95 °C, for up to 30 min. It is desirable to add the xylanase during the feed pelleting process. This would save the feed formulators the additional step of adding liquid xylanase, which is inconvenient and can introduce microbial contamination into the feed. The option of adding solid xylanase as a separate step is also undesirable, as the solids would not be evenly mixed. Marquardt and Bedford (1997. *Enzymes in Poultry and Swine Nutrition*, Marquardt R.R. and Han Z. eds., pp.129-138) indicate that even though currently available enzymes are beneficial for use as feed additives, new enzymes exhibiting high activity and resistance to heat treatment are also desired, however, they note that enzymes exhibiting these properties are not available.

[0005] Xylanases of Family 11 (also termed Family G xylanases) have several properties suitable for feed applications due to their small size and high activity. An example of a moderate temperature Family 11 xylanase is TrX, which is obtained from *Trichoderma reesei*. Moderate temperature xylanases are proven feed additive enzymes with temperature and pH optima compatible with the physiological conditions in the digestive system of animals. However, these enzymes can not tolerate the high temperature of the pelleting process and become inactive during this step.

[0006] Xylanases from high temperature microorganisms (eg. a thermophile), for example *Thermomonospora fusca* xylanase (termed TtX, also a Family 11 xylanase), have also been considered for feed pelleting. The thermostability of such enzymes is sufficient to tolerate the pelleting temperatures. However, thermophilic xylanases have optimum activity at high temperatures (70-80°C), and several of these enzymes have a high pH optimum of 7-9. When introduced into the digesting system of an animal, with a physiological temperature of around 40°C (e.g. poultry 43°C, a similar temperature is noted within swine) and pH of 3-5 in the digesta, these enzymes function poorly.

[0007] Family 11 xylanases have been modified by protein engineering to improve the properties of these enzymes for industrial applications. These modifications have been directed at increasing the temperature and pH optima, along with the thermostability, of these enzymes for specific applications. For example, US 5,405,769 (WO 94/24270) is directed to site-specific mutagenesis of *Bacillus circulans* xylanase (BcX) for the improvement of the thermostability of this enzyme. The disclosed modifications relate to the formation of intermolecular and intramolecular disulfide bonds within BcX, and these modifications resulted in increased thermostability. For example, an improvement in thermostability of up to 6°C with the addition of a single disulfide bond, and up to 10°C with two disulphide bonds was observed. Other modifications included linking the N- and C- termini which increased thermostability by 6°C, or N-terminal mutations, which increased thermostability by 2°C. However, with all of the above modifications the resultant enzymes were either less active (up to 45% less active), or exhibited an increase in the temperature and pH optima. As such these enzymes are not suitable for feed pelleting applications.

[0008] US 5,759,840 also discloses modifications to BcX and *Trichoderma reesei* xylanase (TrX) to increase the

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thermostability; while at the same time increase the temperature and pH optima of these enzymes. Again, these xylanases would not be suitable for feed pelleting applications.

[0009] The above results are in agreement with other reports that note that disulfide bonds are not among the thermostabilization mechanisms employed by thermophilic enzymes (Cowan, D.A., 1995, *Essays Biochem.* 29:193-207), as the disulfide can be broken into dehydroalanine and thioalanine at temperatures over 80°C. Therefore, the enhancement of stability of an enzyme using disulfide bonds is limited to lower temperature ranges. The disulfide bond is thus not recommended to improve the stability of the enzyme at high temperatures (Gupta, M. N., 1991, *Biotech. Applied Biochem.* 14:1-11; Cowan, D.A., 1995, *Essays Biochem.* 29:193-207.).

[0010] None of the above documents address methods for obtaining xylanase enzymes using conventional screening techniques, or by modifying xylanase enzymes, that exhibit the properties of higher temperature tolerance while maintaining optimal performance under conditions of physiological pH and temperature.

[0011] An improvement in the thermostability of *Trichoderma reesei* xylanase II was reported by Paloheimo et al (Paloheimo, M., Mantyla, A., Vehmaanpera, J., Hakola, S., Lantto, R., Lahtinen, T., Parkkinen, E., Fagerstrom, R. and Suominen, P. 1997, in *Carbohydrases from Trichoderma reesei and Other Microorganisms* p255-264). Of the five mutants characterized, the most improved mutant (glutamic acid-38 TrX) retained 50 % of activity at 57°C after 9 min, as compared to 7 min by wild type TrX. Arase et al (Arase, A., Yomo, T., Urabe, I., Hata, Y., Katsube, Y. and Okada, H., 1993, *FEBS Lett.* 316:123-127) describes several modifications to improve the thermostability of a *Bacillus pumilus* xylanase (BpX), however only up to 40% of the residual enzymatic activity was maintained following incubation of these enzymes at a temperature of 57°C for 20 min. Even though, in both of these studies the effects of increased thermostability on pH and temperature optima of the enzymes were not determined, these enzymes exhibit inadequate thermostability for feed pelleting applications.

[0012] In spite of a wide range of experience in screening, testing and modifying xylanase enzymes, there are no reports of xylanases that exhibit the combination of properties required for feed pelleting applications: high thermostability, with optimum activity at physiological pH and temperature. No natural xylanases have been selected, nor has any mutation methodology for the Family 11 xylanases been developed to increase thermostability of xylanase enzymes to, without any change in the temperature and pH optima, and a concomitant loss of the specific activity of the enzyme. Such selected natural xylanases, or xylanases prepared using mutation methodology would offer the advantage of enhancement of feed digestibility and processing in pelleting.

[0013] The present invention is directed to obtaining xylanase enzymes that exhibit the property of increased thermostability, while maintaining the pH and temperature optima that are typically found under physiological conditions.

[0014] It is an object of the invention to overcome disadvantages of the prior art.

[0015] The above object is met by the combinations of features of the main claims, the sub-claims disclose further advantageous embodiments of the invention.

SUMMARY OF THE INVENTION

[0016] The present invention relates to thermostable xylanase enzymes. More specifically, the present invention is directed to thermostable xylanase enzymes that exhibit high activity at or near physiological pH and temperature, and the use of these xylanase enzymes in feed pelleting applications.

[0017] According to the present invention there is provided an isolated Family 11 xylanase, comprising at least one intramolecular disulfide bond, and a basic amino acid at position 162 (*Trichoderma reesei* xylanase II numbering) or its equivalent, said position determined from sequence alignment of said isolated xylanase with the *Trichoderma reesei* xylanase II amino acid sequence defined in SEQ ID NO:18, said isolated xylanase exhibiting at least 40% of optimal activity from about pH 3.5 to about pH 6.0, and from about 40 to about 60°C, and at least 30% of optimal activity after a pre-incubation step for 30 minutes at 70°C, 80°C or 90°C in the presence of 40% glycerol; a pre-incubation step for 30 or 60 minutes at 62.5°C in the absence of a stabilizer; or a pre-incubation step of 30 minutes at 64°C or 66°C in the absence of a stabilizer. The basic amino acid is selected from the group consisting of lysine, arginine and histidine. Preferably, the basic amino acid is histidine.

[0018] The modified xylanase of the invention comprises at least one disulfide bridge. Preferably, the modified xylanase comprises one or two disulfide bridges.

[0019] The present invention is directed to an isolated Family 11 xylanase. Furthermore, this invention pertains to an isolated xylanase, wherein the Family 11 xylanase is from *Trichoderma*.

[0020] The present invention is also directed to the isolated xylanase as defined above wherein said xylanase is selected from the group consisting of TrX-162H-DS1, TrX-162H-DS2, and TrX-162H-DS4.

[0021] This invention also includes a method of obtaining a Family 11 xylanase comprising:

- i) selecting an organism that expresses xylanase activity, and obtaining said xylanase from the organism;
- ii) determining whether the xylanase exhibits at least 40% of optimal activity from pH 3.5 to pH 6.0, and from 40 to

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60°C; and

iii) determining whether the xylanase is a Family 11 xylanase and whether the xylanase exhibits at least 30% of optimal activity after a pre-incubation step for 30 minutes at 70°C, 80°C or 90°C in the presence of 40% glycerol; a pre-incubation step for 60 minutes at 62.5°C in the absence of a stabilizer; or a pre-incubation step of 30 minutes at 64°C or 68°C in the absence of a stabilizer;

iv) determining whether said xylanase has a basic amino acid at position 162 (*Trichoderma reesei* xylanase II numbering); said position determined from sequence alignment of said isolated xylanase with *Trichoderma reesei* xylanase II amino acid sequence defined in SEQ ID NO:16, and an intra-molecular disulfide bond; and

v) retaining the xylanase that expresses these properties.

Step i) of the above method may also include partially purifying the xylanase.

[0022] The present invention also pertains to a method of preparing animal feed, wherein the method comprises applying the isolated, modified, Family 11 xylanase as defined above onto the animal feed to produce a xylanase-animal feed combination, and heat sterilizing the xylanase-animal feed combination. Preferably, the animal feed is a poultry or swine feed.

[0023] The present invention also pertains to an isolated Family 11 xylanase characterized in comprising at least one intramolecular disulfide bond, and a basic amino acid at position 162 (*Trichoderma reesei* xylanase II numbering) or its equivalent, said position determined from sequence alignment of said isolated xylanase with the *Trichoderma reesei* xylanase II amino acid sequence defined in SEQ ID NO: 16, said xylanase exhibiting at least 30% of optimal activity after a pre-incubation step for 30 minutes at 70°C, 80°C or 90°C in the presence of 40% glycerol; a pre-incubation step for 30 or 60 minutes at 62.5°C in the absence of a stabilizer; or a pre-incubation step of 30 minutes at 64°C or 68°C in the absence of a stabilizer.

[0024] The present invention is directed to obtaining xylanase enzymes that exhibit pH and temperature optima that are found within the digesta of an animal, while at the same time the xylanase molecule exhibits thermostability and can therefore withstand processes associated with sterilizing and producing pelleted feed. The prior art discloses obtaining thermostable enzymes, either through selection of native enzymes or through genetic engineering, however, these enzymes do not exhibit physiological pH and temperature optima. The prior art also discloses xylanase enzymes that exhibit optimal enzyme activity at physiological pH and temperature, however, these enzymes are not thermally stable. Furthermore, there is nothing in the prior art to suggest that native xylanase enzymes, or that xylanase enzymes may be modified as disclosed herein in order to obtain xylanase enzymes that exhibit high temperature tolerance suitable for feed pelleting, and retain optimum enzymatic activity at or near physiological conditions.

[0025] This summary of the invention does not necessarily describe all necessary features of the invention but that the invention may also reside in a sub-combination of the described features.

BRIEF DESCRIPTION OF THE DRAWINGS

[0026] These and other features of the invention will become more apparent from the following description in which reference is made to the appended drawings wherein:

FIGURE 1 shows the multiple amino acid sequence alignment among Family 11 xylanases. The amino acids common to at least 80% of the Family 11 xylanases listed are indicated in bold. The residues common to all Family 11 xylanases are underlined. *Bacillus pumilus* (Bp); *Clostridium acetobutylicum* P262 XynB (Ca); *Clostridium stercoararium* (Cs); *Ruminococcus flavefaciens* (Rf); *Trichoderma reesei* XynII (Tr2); *Trichoderma viride* (Tv); *Trichoderma harzianum* (Th); *Schizophyllum commune* Xylanase A (Sc); *Aspergillus niger* var. awamori (An); *Aspergillus tubigensis* (At); *Trichoderma reesei* XynI (Tr1); *Streptomyces* sp. No. 36a (Ss); *Streptomyces lividans* Xylanase B (S1B); *Streptomyces lividans* Xln C (S1C); *Thermomonospora fusca* TrxA (Tf); *Bacillus circulans* (Bc); *Bacillus subtilis* (Bs)

FIGURE 2 shows the synthetic oligonucleotides for the construction of gene sequence encoding the *Trichoderma* xylanase in the plasmid pTrX (SEQ ID NO:18).

FIGURE 3 shows the effect of incubation time on the residual enzymatic activity of mutant TrX, TrX-DS1, TrX-162H, TrX-162H-DS1, and TrX-162H-DS4 at 62.5°C. The data are normalized to that observed at 0 min.

FIGURE 4 shows the effect of temperatures on the residual enzymatic activity of several of the modified xylanases of the present invention. Figure 4(a) shows the residual enzymatic activity of TrX, TrX-DS1, TrX-162H-DS1, TrX-162H-DS2, and TrX-162H-DS4 in sodium citrate buffer in a 30 min incubation. Figure 4(b) shows the effect of temperatures on the residual enzymatic activity of the mutant TrX-DS8. For Figures 4(a) and

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(b) The data are normalized to that observed at 48°C. The T_{50} , which is the incubation temperature allowing the maintenance of 50% residual activity after 30 min, was determined for each mutant TrX.

5 **FIGURE 5** shows the effect of temperatures on the residual enzymatic activity of mutant TrX, TrX-DS1 and TrX-162H-DS1 in 40% glycerol in a 30 min incubation. The data are normalized to that observed at 50°C.

FIGURE 6 shows the effect of incubation time on the residual enzymatic activity of TrX-162H-DS1 in 40% glycerol at 90°C. The data are normalized to that observed at 0 min.

10 **FIGURE 7** shows the effect of temperature on release of xylose in a 30 min hydrolysis of soluble xylan by TrX, TrX-162H-DS1, TrX-162H-DS2 and TrX-162H-DS4 at pH 4.5. The data are normalized to that observed at the temperature optimum.

15 **FIGURE 8** shows the effect of pH on the release of xylose in a 7 min hydrolysis of soluble xylan by TrX, TrX-162H-DS1, TrX-162H-DS2 and TrX-162H-DS4 at 40°C. The data are normalized to that observed at the pH optimum.

DESCRIPTION OF PREFERRED EMBODIMENT

20 **[0027]** The present invention relates to thermostable xylanase enzymes and their use as feed additives. More specifically, the present invention is directed to thermostable xylanase enzymes that show good thermostability and exhibit high activity at or near physiological pH and temperature.

[0028] The following description is of a preferred embodiment by way of example only and without limitation to the combination of features necessary for carrying the invention into effect.

25 **[0029]** By physiological pH and temperature, it is meant the range in temperature and pH compatible with the digestive system within an animal, for example but not limited to, poultry and swine. For example, a suitable physiological temperature range is from about 35 to about 60°C, more preferably, this range is from about 40 to about 50°C. Similarly, a suitable physiological pH range is from about pH 3.0 to about 7.0, preferably, this range is from about pH 3.5 to about 6.0. The time required for the digestion of feed within the gut of an animal varies from animal to animal. For example, in swine digestion of feed is from about 2 to about 4 hours, while in poultry it is up to about 12 hours.

30 **[0030]** By high activity at physiological pH and temperature, it is meant that the enzyme exhibits at least 40% of its optimum activity at physiological pH and temperature. The optimum pH and temperature-range can be outside the physiological range, provided that the enzyme exhibits at least 40% of its optimum activity within the physiological range, for example from about 40 to about 50°C and pH from about 3.5 to about 6. Examples 4 and 5 describe the determination of a suitable xylanase enzyme that exhibits these properties.

35 **[0031]** "Thermostable" or "thermostability" as used herein refer to a property of an enzyme. An enzyme is considered to be thermostable if it exhibits at least one of the following properties:

40 1) the enzyme exhibits at least 30% of its optimal activity following a pre-incubation step of 30 min at 70°C, 80°C, or 90°C, at pH 5.0, in the presence of a stabilizing agent such as 40% glycerol. Preferably, the enzyme exhibits at least 40% of its optimal activity following a 30 min, 70°C pre-incubation step in glycerol, for example but not limited to, TrX-162H-DS1 (Figure 5);

45 2) the enzyme exhibits 30% of its optimal activity following a pre-incubation step of 30 or 60 min at 62.5°C in the absence of a stabilizer. Preferably, the enzyme exhibits at least 40% of its optimal activity following a 30 min pre-incubation, for example but not limited to, TrX-162H-DS1 and TrX-162H-DS4 (Figure 3);

50 3) the enzyme exhibits at least 30% of its optimal activity following a preincubation step of 30 min at 64°C in the absence of a stabilizer. Preferably, the enzyme exhibits at least 40% of its optimal activity following the 30 min, 64°C pre-incubation step, for example but not limited to, TrX-162H-DS1 and TrX-162H-DS4 (Figure 4); or

55 4) the enzyme exhibits at least 30% of its optimal activity following a preincubation step of 30 min at 68°C in the absence of a stabilizer. Preferably, the enzyme exhibits at least 40% of its optimal activity following the 30 min, 68°C pre-incubation step, for example but not limited to, TrX-162H-DS1 and TrX-162H-DS4 (Figure 4). In each of the above cases, the optimal activity of the enzyme is determined at an optimal pH and temperature for that enzyme in the presence or absence of stabilizer as required.

[0032] By "TrX numbering" it is meant the numbering associated with the position of amino acids based on the amino

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acid sequence of TrX (Xyn II - Table 1; Tr2 - Figure 1). As disclosed below and as is evident upon review of Figure 1, Family 11 xylanases exhibit a substantial degree of sequence homology. Therefore, by aligning the amino acids to optimize the sequence similarity between xylanase enzymes and by using the amino acid numbering of TrX as the basis for numbering, the positions of amino acids within other xylanase enzymes can be determined relative to TrX.

[0033] By modified xylanase, it is meant the alteration of a xylanase molecule using techniques that are known to one of skill in the art. These techniques include, but are not limited to, site directed mutagenesis, cassette mutagenesis, synthetic oligonucleotide construction, cloning and other genetic engineering techniques. Alterations of a xylanase enzyme, in order to produce a modified xylanase may also arise as a result of applying techniques directed at inducing mutations within native or genetically engineered xylanases via the addition of known chemical mutagens, UV exposure, or other treatments known to induce mutagenesis within a host organisms that express a xylanase of interest. Such techniques are well known within the art.

[0034] Table 1 lists the Family 11 xylanases free of cellulase activity. These enzymes share extensive amino acid sequence similarity and possess amino acids common to Family 11, for example two glutamic acid (E) residues serving as the essential catalytic residues, amino acids 86 and 177 (using TrX numbering). Structural comparisons of several Family 11 xylanases via X-ray crystallography indicates that these Family 11 xylanases of bacterial and fungal origins share the same general molecular structure (see for example US 5,405,769; Arase, A., Yomo, T., Urabe, I., Hata, Y., Katsube, Y. and Okada, H, 1993, FEBS Lett. 316:123-127). Most of the family 11 xylanases identified so far are mesophilic and have low-molecular mass (20kDa).

TABLE 1: Family 11 xylanases

Microbe	Xylanase	Ref. In Figure 1	Sequence Listing
<i>Aspergillus niger</i>	Xyn A	An	SEQ ID NO: 1
<i>Aspergillus kawachii</i>	Xyn C		
<i>Aspergillus tubingensis</i>	Xyn A	A1	SEQ ID NO: 2
<i>Bacillus circulans</i>	Xyn A	Bc	SEQ ID NO: 3
<i>Bacillus pumilus</i>	Xyn A	Bp	SEQ ID NO: 4
<i>Bacillus subtilis</i>	Xyn A	Bs	SEQ ID NO: 5
<i>Cellulomonas fimi</i>	Xyn D		
<i>Chaetia spp.</i>	Xyn		
<i>Clostridium acetobutylicum</i>	Xyn B	Ca	SEQ ID NO: 6
<i>Clostridium stercoarum</i>	Xyn A	Cs	SEQ ID NO: 7
<i>Fibrobacter succinognees</i>	Xyn C		
<i>Neocallimastix patriciarum</i>	Xyn A		
<i>Nocardopsis dassonvillei</i>	Xyn II		
<i>Ruminococcus flavefaciens</i>	Xyn A	Rf	SEQ ID NO: 8
<i>Schizophyllum commune</i>	Xyn	Sc	SEQ ID NO: 9
<i>Streptomyces lividans</i>	Xyn B	S1B	SEQ ID NO: 10
<i>Streptomyces lividans</i>	XynC	S1C	SEQ ID NO: 11
<i>Streptomyces sp. No. 36a</i>	Xyn	Ss	SEQ ID NO: 12
<i>Streptomyces thermoviolaceus</i>	XynII		
<i>Thermomonospora fusca</i>	Xyn A	Tf	SEQ ID NO: 13
<i>Trichoderma harzianum</i>	Xyn	Th	SEQ ID NO: 14
<i>Trichoderma reesei</i>	Xyn I	Tr1	SEQ ID NO: 15
<i>Trichoderma reesei</i>	Xyn II	Tr2	SEQ ID NO: 16
<i>Trichoderma viride</i>	Xyn	Tv	SEQ ID NO: 17

[0035] It is considered within the scope of the present invention that xylanases, including Family 11 xylanases for example but not limited to *Trichoderma reesei* xylanase II, *Trichoderma reesei* xylanase I, *Trichoderma viride* xylanase, *Streptomyces lividans* xylanase B and *Streptomyces lividans* xylanase C, may be modified following the general approach and methodology as outlined herein. It is also considered within the scope of the present invention that non-Family 11 xylanases may also be modified following the general principles as described herein in order to obtain a xylanase enzyme that is thermostable and exhibits high activity at physiological pH and temperature.

[0036] Furthermore, native xylanases may also be obtained by using standard screening protocols in order to identify enzymes that exhibit the properties of increased thermostability yet maintaining high activity at physiological temperature

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and pH. Such protocols involve:

- selecting of a desired organism, for example a thermophile;
- extracting or obtaining the xylanase from the organism, and partially purifying the enzyme if desired; and
- characterizing the extracted enzyme to determine whether the enzyme is thermostable, as defined above (in the presence or absence of a stabilizing agent, such as glycerol), determining the enzymes pH and temperature optima, and determining the activity of the enzyme at physiological pH and temperature.

[0037] Any enzymes identified using the above protocol that exhibit thermostability and high activity at physiological pH and temperature may be used as animal feeds.

[0038] The present invention also relates to modified xylanase enzymes that exhibit increased thermostability while maintaining high activity at physiological pH and temperature. For example, and without wishing to limit the present invention in any manner, a modified *Trichoderma reesei* xylanase (TrX) is disclosed that exhibits increased thermostability while maintaining pH and temperature optima at or near physiological range. Two modifications in the TrX were combined in order to obtain a novel xylanase (TrX-162H-DS1). The first modification includes a double mutation to create two cysteines for the formation of a single disulfide bond. Such a modification has been described for *Bacillus circulans* xylanase (C100/C148; BcX amino acid numbering) in US 5,405,769. However, this mutation bestows only a minor increase in the ability of the enzyme to withstand high temperatures (see TrX-DS1, Figures 3-5) and this modification is not adequate to produce an enzyme capable of surviving high temperatures associated with the pelleting process. When this mutation is combined with a second mutation as per the teaching of this invention, involving the substitution of a basic amino acid such as histidine (H) for glutamine (Q) in position 162, the resultant combination mutant xylanase exhibits the desired properties of thermostability (TrX-162H-DS1; see Figures 5 and 6), and greater than 40% of optimum activity at physiological pH (Figure 8), and temperature (Figure 7).

[0039] Another mutant xylanase in the present invention, TrX-162H-DS4 differs from TrX-162H-DS1 by possessing an additional disulfide (108/158, that is between positions 108 and 158). This type of double disulfide mutant has previously been described for the xylanase of *Bacillus circulans* (C98/C152, 100/148; BcX amino acid numbering; Wakarchuck et al., 1994 Protein Engineering, 7:1379-1386). The BcX mutant does not comprise an equivalent basic amino acid (e.g. H for Q at position 162) substitution as disclosed herein. The mutant TrX-162H-DS4 shows a dramatic increase of thermostability (see Figure 4(a)), with an increase in the T_{50} of TrX-162H-DS4 of 14°C. This is an improvement over the prior art double disulfide BcX mutant which exhibits an increase in the T_{50} of 10°C, thereby demonstrating the contribution of the Q162H mutation in the disulfide mutants of TrX.

[0040] The present invention also pertains to additional mutations that have been found to be effective in producing a xylanase that exhibits thermostability and a desirable pH profile. An example of such mutations may be found in, but are not limited to, TrX-DS8. TrX-DS8 includes the mutations listed for N1-TX13 as disclosed in US 5,759,840, namely N1OH, Y27M and N29L, and also includes N44D, Q125A, I129E, Q162H and a disulfide bond between positions 110 and 154. TrX-DS8 exhibits the property of thermostability (Figure 4(b)), a pH profile paralleling that of TrX-162-DS1, and greater than 40 % of optimum activity at physiological pH, and temperature.

[0041] Xylanase enzymes comprising the substitution of H for Q at position 162 (termed Q162H) in isolation has been reported in US 5,759,840, however, these mutants exhibited no improvement in thermostability or other properties over natural TrX. However, by combining these two modifications, several novel xylanases (TrX-162H-DS1, TrX-162H-DS2 and TrX-162H-DS4) were obtained with improved thermostability. This property was not observed with either mutation alone. Furthermore, these modified xylanases exhibit high activity at or near physiological temperature and pH. These mutations are also found in TrX-DS8, which also exhibits improved thermostability and high activity at or near physiological conditions.

[0042] Following the methods of the present invention novel xylanase enzymes may be obtained that are far more suitable for feed pelleting applications than enzymes currently available. Similar modifications may be made in other Family 11 xylanases, including but not limited to, xylanase enzymes obtained from *Trichoderma*, *Streptomyces* and *Schizophyllum*. However, it is also within the scope of the present invention that other xylanase enzymes, in addition to Family 11 xylanases can be modified as disclosed herein in order to obtain xylanases with that are thermostable and exhibit high activity at physiological pH and temperature. Furthermore, it is within the scope of the present invention that native xylanase enzymes with the properties of thermostability and high activity at physiological pH and temperature may be obtained following screening protocols that select for both thermostability and high activity at physiological pH and temperature.

[0043] In use, the formulation of the feed enzyme can improve the enzymes thermostability, as adsorption into feed improves stability as the enzyme is brought into contact with its substrate. Therefore, in determining thermostability of the xylanases of the present invention, xylanases were characterized in the presence and absence of stabilizing agents, for example but not limited to glycerol. Fisk and Simpson (1993) have reported that 40% glycerol enhanced the temperature tolerance of wild type TrX by less than + 10°C, however, this is much less stability than the enzymes of the present

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invention. The combination-mutant xylanases of the present invention can tolerate incubation in buffer at a higher temperature (59-69°C), as compared to natural xylanase (55°C; also see Figure 3 and 4). In the presence of 40% glycerol, the combination mutants can retain a substantial portion of their activity at 70 to 90°C (see figure 5), while the natural xylanase is totally inactivated at these temperatures..

[0044] One of the modifications to the combination mutant xylanase as proposed herein is the substitution of amino acid 162 (TrX numbering, based on Tr2 in Figure 1; which for TrX is glutamine) with the basic amino acid histidine (termed Q162H). However, it is considered within the scope of the present invention that other amino acids may also be substituted at this position. Preferably the substituted amino acid is basic (positively charged), for example lysine (Q162K) or arginine (Q162R). It has been observed herein that the substitution at the position 162, or its equivalent in other Family 11 xylanases, by a basic amino acid such as histidine can greatly improve the thermostability of a xylanase enzyme that comprises at least one intramolecular disulfide bond. Importantly, it has also been observed herein that this substitution at position 162 not only increases thermostability but also does not significantly change the temperature and pH profiles, and the specific activity of the modified xylanase.

[0045] Histidine-162 residue (TrX numbering) in the combination mutant is found in several natural Family 11 xylanases, such as those of *Trichoderma harzianum*, *Aspergillus niger*, var. *awamori*, *Aspergillus tubingensis*, *Thermomonospora fusca*, *Bacillus circulans* and *Bacillus subtilis* in the corresponding position. Similarly, *Clostridium acetobutylicum* comprises a lysine at this equivalent position. However, all, of these xylanases, with the exception of the *Thermomonospora fusca* xylanase, are produced by mesophilic hosts and exhibit low thermostability. As a result there is no evidence to suggest any beneficial effect on thermostability by presence of a basic amino acid residue at this position. In the *Thermomonospora fusca* xylanase, the N-terminal sequence (1-29) which is distant from the site of the present invention, has been shown to contribute to thermostability, and there is no evidence to suggest that thermostability may be associated with a histidine at this equivalent position (i.e. TrX 162).

[0046] This invention is also directed to xylanases that comprise at least one modification that results in increased thermostability while maintaining high activity at physiological pH and temperature. For example, native *Schizophyllum commune* xylanase has a disulfide bond at positions 110/154 (TrX numbering). However, this enzyme exhibits low thermostability. Therefore, this enzyme can be modified using the methods of the present invention to substitute a basic amino acid, either histidine, arginine or lysine for the naturally occurring leucine at position 200 of *Schizophyllum commune* (which is equivalent to position 162 using TrX numbering; see Figure 1; Sc). Therefore, increased thermostability can be achieved through a one-step modification.

[0047] Also considered within the scope of the present invention are combination mutants comprising both an intramolecular disulfide bond and a basic amino acid substitution as outlined above. The intramolecular disulfide bond may arise as a result of a mutation at one or more specific residues, for example (per TrX numbering):

- residues-110/-154, for example, but not limited to TrX-162H-DS1 or TrX-DS8;
- residues-108/-158, for example, but not limited to TrX-162H-DS2; or
- residues-108/-158, -110/-154, for example, but not limited to TrX-162H-DS4.

[0048] Also considered within the scope of the present invention are modifications of thermostable xylanases, for example, but not limited to TrX. These modifications maintain the thermostability of the native enzyme, yet alter the pH and temperature optima so that they exhibit high activity at physiological pH and temperature not normally associated with the enzyme.

TABLE 2: Modified xylanases

XYLANASE	DESCRIPTION
wild type TrX	wild type <i>T. reesei</i> xylanase.
TrX-162H	TrX mutant with mutation Q162H.
TrX-DS1	TrX mutant with an intramolecular disulfide bond between positions-110 and 154.
TrX-162H-DS1	TrX mutant with two mutations, (i) a disulfide bond between positions-110 and 154, and (ii) mutation Q162H.
TrX-162H-DS2	TrX mutant with two mutations, (i) an intramolecular disulfide bond between positions-108 and 158, and (ii) mutation Q162H.
TrX-162H-DS4	TrX mutant with two mutations, (i) two intramolecular disulfide bonds at residues-110/154 and residues-108/158, and (ii) mutation Q162H.

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(continued)

XYLANASE	DESCRIPTION
TrX-DS8	Trx mutant with i) an intramolecular disulfide bond between positions-110 and 154, and ii) comprising mutations N10H, Y27M, N29L, N44D, Q125A, 1129E, and Q162H

[0049] The present invention will be further illustrated in the following examples. However it is to be understood that these examples are for illustrative purposes only, and should not be used to limit the scope of the present invention in any manner.

Examples:

Example 1: Construction of the *Trichoderma reesei* mutant xylanases

[0050] Basic recombinant DNA methods like plasmid preparation, restriction enzyme digestion, polymerase chain reaction, oligonucleotide phosphorylation, ligation, transformation and DNA hybridization were performed according to well-established protocols familiar to those skilled in the art (Sung, W. L., Yao, F.-L., Zahab, D. M. and Narang, S. A. (1986) Proc. Natl. Acad. Sci. USA 83:561-565) or as recommended by the manufacturer of the enzymes or kit. The buffer for many enzymes have been supplied as part of a kit or constituted following to the instruction of the manufacturers. Restriction enzymes, T4 polynucleotide kinase and T4 DNA ligase were purchased from New England BioLabs LTD, Mississauga, Ont. A precursor plasmid pXYbc has previously prepared and published (Sung, W. L., Luk, C. K., Zahab, D. M. and Wakarchuk, W. (1993) Protein Expression Purif. 4:200-208; US 5,405,769). A commonly used *E. coli* strain, HB101 (clonotech Lab, Palo Alto, CA) was used as transformation and expression host for all gene construct. Birchwood xylan was purchased from Sigma (St. Louis, Mo). Hydroxybenzoic acid hydrazide (HBAH) was purchased from Aldrich. Oligonucleotides were prepared with an Applied Biosystem DNA synthesizer, model 380B. Xylanase assays have been performed in a covered circulating water bath (Haake type F 4391) with a fluctuation of $\pm 0.1^{\circ}\text{C}$. Temperature of the water bath was confirmed with a thermocouple.

A. Construction of the precursor plasmid pTrX

[0051] The precursor plasmid pTrX for all subsequent mutations is published (Sung et al, 1995). This plasmid is derived from a pUC119 plasmid with a synthetic nucleotide sequence encoding a *Trichoderma reesei* xylanase inserted (Figure 2). Expression of this xylanase and other mutant xylanases subsequently described are under the control of the lac promoter of the pUC plasmid. The total assembly of the gene required two stages, initially for the (92-190) region, then followed by the (1-92) region. The protocol for the construction of this gene is routine and identical to the standard published procedure for many other genes. It required enzymatic phosphorylation of overlapping synthetic oligonucleotides which encodes xylanase. This was followed by their ligation into a appropriately cut plasmid pUC119.

[0052] Initially ten overlapping oligonucleotides:

XyTv-101,	SEQ ID NO:28
XyTv-102,	SEQ ID NO:29
TrX-103,	SEQ ID NO:30
XyTv-104,	SEQ ID NO:31
XyTv-105,	SEQ ID NO:32
XyTv-106,	SEQ ID NO:33
XyTv-107,	SEQ ID NO:34
TrX-108,	SEQ ID NO:35
XyTv-109,	SEQ ID NO:22
XyTv-110,	SEQ ID NO:36

encoding the TrX(92-190) sequence (Figure 2), were designed with codon usage frequency imitating that of *E. coli* (Chen et al. 1982). The SalI and BglII cohesive ends of two terminal oligonucleotides enabled the enzymatic ligation of the ten fragments to the linearized plasmid pXYbc. The ten oligonucleotides (50 pmol, 1 L for each) encoding the TrX(92-190) was phosphorylated in a mixture containing 10X standard kinase buffer (0.4 L), 1mM ATP (4 L), T4 DNA kinase (5 units), and water (3 L). Phosphorylation reaction was carried out for 1 h at 37°C . The solutions were then combined and heated to 70°C for 10 min. After being cooled slowly to room temperature, the combined solutions were added to a mixture of

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4mM ATP (3.5 L), EcoR1-HindIII linearized plasmid pUC119 (0.1 pmol), and T4 DNA ligase (3.5 L) and incubated at 12°C for 20 h. Aliquots of the ligation mixture were used to transform *E. coli* HB101 in YT plate (8 g yeast extract, 5 g bacto-tryptone, 5 g NaCl, 15 g of agar in 1 L of water) containing ampicillin (100 mg/L).

[0053] For the preparation of a hybridization probe, one of the oligonucleotide XyTv-110 (10 pmol, 1 L) was phosphorylated ³²P-ATP (10 pmol, 3 L) in T4 DNA kinase (1 L), 10X kinase buffer (1 L), and water (4 L) at 37°C for 1 h.

[0054] Transformants were selected randomly for hybridization analysis. Colonies were grown on nylon filters on YT plates with ampicillin overnight. They were then denatured with 0.5N NaOH -1.5M NaCl (10 min) and neutralized with 0.5N Tris-HCl (pH 7.0) - 1.5M NaCl (10 min). After irradiation by UV of 254 nm for 8 min, the filters were washed with 6X SSC - 0.05% Triton X-100 for 30 min. Cell debris was scraped off completely. After another 30 min. in fresh solution, the duplicate filters were transferred individually into separate mixtures of 6X SSC - 1 % dextran sulphate - 0.05 % TritonX-100 - 1X Denhardt's hybridization fluid. The ³²P-labelled probe was added to the filter. After 16 h at 45°C, the filter was washed twice with 6X SSC - 0.05 % TritonX-100 at room temperature for 5 min. and then at 65°C for 30 min. Positively hybridized clones with the intermediate plasmid pBcX.TrX were identified by auto-radiographic analysis.

[0055] The above protocol, involving enzymatic phosphorylation of synthetic overlapping oligonucleotides and ligation into a linearized plasmid, has again been used in the assembly of the TrX(1-92) region and in the cassette mutagenesis for the subsequent generation of other mutant series described in this invention.

[0056] For the assembly of the TrX(1-92) region to complete the full-length *Trichoderma* gene, the intermediate plasmid pBcX.TrX was linearized by NheI and KpnI endonucleases to release the DNA Insert for BcX(1-83). With NheI and KpnI cohesive ends, eight overlapping oligonucleotides:

TrX-1,	SEQ ID NO:37
XyTv-2,	SEQ ID NO:38
TrX-3,	SEQ ID NO:39
XyTv-4,	SEQ ID NO:40
XyTv-5,	SEQ ID NO:41
TrX-6,	SEQ ID NO:42
XyTv-7,	SEQ ID NO:43
TrX-8,	SEQ ID NO:44,

encoding the published TrX(1-91) sequence were ligated into the linearized plasmid pBcX.TrX (Figure 2), via the protocol described above. The new plasmid pTrX therefore harbored a synthetic TrX gene (SEQ ID NO: 18).

[0057] All mutant xylanases described below have been constructed via the method of cassette mutagenesis as described above. The protocol for the cassette mutagenesis was identical to that for gene assembly fully described above. Such cassette mutagenesis involved (i) enzymatic phosphorylation of overlapping synthetic oligonucleotides, (ii) their ligation with the linearized plasmid, (iii) transformation into the *E. coli* HB101 competent cells, (iv) identification of the mutant transformants via hybridization with the labelled oligonucleotide as probe, and (v) confirmation of the mutation through dideoxy nucleotide sequencing.

B. Construction of the plasmid pTrX-DS1

[0058] The mutant TrX-DS1 (SEQ ID NO's:54, 55) was identical to TrX with a covalent disulfide bond between residues-110 and 154. This was accomplished through two single mutations, ie. conversion of both residues serine-110 and asparagine-154 to cysteine. Upon expression of the mutant xylanase, these two cysteine residues will form a disulfide bond. The construction of the plasmid pTrX-DS1 was through ligation of the following overlapping phosphorylated oligonucleotides:

TX-110C	SEQ ID NO: 19,
TX-110C-2	SEQ ID NO:20,
TX-103b	SEQ ID NO:21,
XyTv-109	SEQ ID NO:22,
TX-108b	SEQ ID NO:23,
TX-154C	SEQ ID NO:24,
TX-154C-2	SEQ ID NO:25,

into KasI/AvrII-linearized plasmid pTrX in a cassette mutagenesis as shown below.

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TX-110C-2

101 102 103 104 105 106 107 108 109 110 111 112 113 114 115
 G A T K L G E V T C D G S V Y
 5'-GC GCC ACA AAA TTA GGC GAA GTC ACT TGT GAT GGA TCC GTA TAT
 3'-G TGT TTT AAT CCG CTT CAG TGA ACA CTA CCT AGG CAT ATA
 Kasi| TX-110C |

TX-103b

116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131
 D I Y R T Q R V N Q P S I I G T
 GAT ATC TAC CGT ACC CAA CGC GTT AAT CAG CCA TCG ATC ATT GGA ACC
 CTA TAG ATG GCA TGG GTT GCG CAA TTA GTC GGT AGC TAG TAA CCT TGG
 XyTV-109

132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147
 A T F Y Q Y W S V R R N H R S S
 GCC ACC TTT TAT CAG TAC TGG AGT GTT AGA CGT AAT CAT CGG AGC TCC
 CGG TGG AAA ATA GTC ATG ACC TCA CAA TCT GCA TTA GTA GCC TCG AGG
 TX-108b

TX-154C-2

148 149 150 151 152 153 154 155 156 157 158 159 160 161 162 163
 G S V N T A C H F N A W A Q Q G
 GGT TCG GTT AAT ACT GCA TGC CAC TTT AAT GCC TGG GCA CAG CAA GGG
 CCA AGC CAA TTA TGA CGT ACG GTG AAA TTA CGG ACC CGT AGT GTT CCC
 SphI TX-154C

164 165 166 167
 L T L G
 TTA ACC
 AAT TGG GAT C

AvrII |

C. Construction of the plasmid pTrX-162H-DS1

[0059] The mutant TrX-162H-DS 1 (SEQ ID NO:56) was identical to TrX-DS 1 with a single mutation of glutamine-162 into histidine. The construction of the plasmid pTrX-162D-DS 1 was through ligation of oligonucleotides:

TX-162H-3 SEQ ID NO: 26, and
 TX-162H-4 SEQ ID NO: 27

into SphI/AvrII-linearized plasmid pTrX-DS1 in a cassette mutagenesis, as shown below.

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TX-162H-3

153 154 155 156 157 158 159 160 161 162 163 164 165 166 167
 A C H F N A W A Q H G L T L G
 5' -C CAC TTC AAT GCA TGG GCA CAG CAC GGG TTA ACC
GT ACC GTG AAG TTA CGT ACC CGT GTC GTG CCC AAT TCG GAT C-5'
 SphI AVRII

TX-162H-4

D. Construction of the plasmid pTrX-162H-DS2

[0060] The mutant TrX-162H-DS2 (SEQ ID NO's:57,58) was identical to TrX, but with a covalent disulfide bond between residues-108 and -158, and a mutation glutamine-162 to histidine. The 108/110 disulfide required two single mutations, ie, conversion of both residues valine-108 and alanine-158 to cysteine. Upon expression of the mutant xylanase, these two cysteine residues will form a disulfide bond. The construction of the plasmid pTrX-162H-DS2 was through ligation of the following overlapping phosphorylated oligonucleotides:

TX-108C	SEQ ID NO:45,
TX-108C-2	SEQ ID NO:46,
TX-103b	SEQ ID NO:21,
XyTv-109	SEQ ID NO:22,
TX-108b	SEQ ID NO:23,
TX-158C-162H	SEQ ID NO:47, and
TX-158C-162H-2	SEQ ID NO:48

Into the KasI/AvrII-linearized plasmid pTrX in a cassette mutagenesis as shown below.

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      |                               TX-108C-2
101 102 103 104 105 106 107 108 109 110 111 112 113 114 115
  G A T K L G E C T S D S S V Y
5 5-GC GCC ACA AAA TTA GGC GAA TGC ACT AGT GAT GGA TCC GTA TAT
   3'-G TGT TTT AAT CCG CTT ACG TGA TCA CTA CCT AGG CAT ATA
      KasI|                               TX-108C      |

116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131
  D I Y R T Q R V N Q P S I I G T
15 GAT ATC TAC CGT ACC CAA CGC GTT AAT CAG CCA TCG ATC ATT GGA ACC
   CTA TAG ATG GCA TGG GTT GCG CAA TTA GTC GGT AGC TAG TAA CCT TGG
      XyTV-109

132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147
  A T F Y Q Y W S V R R N K R S S
25 GCC ACC TTT TAT CAG TAC TGG AGT GTT AGA CGT AAT CAT CGG AGC TCC
   CGG TGG AAA ATA GTC ATG ACC TCA CAA TCT GCA TTA GTA GCC TCG AGG
      |                               TX-108b

30 TX-158C-162H-2

148 149 150 151 152 153 154 155 156 157 158 159 160 161 162 163
  G S V N T A N H F N C W A Q H G
35 GGT TCG GTT AAT ACT GCA AAT CAC TTT AAT TGC TGG GCA CAG CAC GGG
   CCA AGC CAA TTA TGA CGT TTA GTG AAA TTA ACG ACC CGT AGT GTG CCC
      |                               TX-158C-162H

164 165 166 167
  L T L G
40 TTA ACC
   AAT TGG GAT C
45 AvxII |

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E. Construction of the plasmid pTrX-162H-DS4

50 [0061] The mutant TrX-162H-DS4 (SEQ IDNO's:59, 60) was identical to TrX, but with two covalent disulfide bonds 108/158 and 110/154 and a mutation glutamine-162 to histidine. The two disulfides required four single mutations, ie. conversion of the residues valine-108, serine-110, asparagine-154 and alanine-158 to cysteine. Upon expression of the mutant xylanase, these four cysteine residues will form two disulfide bonds. The construction of the plasmid pTrX-162H-DS4 was through ligation of the following overlapping phosphorylated oligonucleotides:

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TX-108C-110C      SEQ ID NO:49,
TX-108C-110C-2    SEQ ID NO:50,

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(continued)

TX-103b SEQ ID NO:21,
XyTv-109 SEQ ID NO:22,
TX-108b SEQ ID NO:23,
TX-154C-158C-162H SEQ ID NO:51 and
TX-154C-158C-162H-2 SEQ ID NO:52

into the Kasi/AvrII-linearized plasmid pTrX In a cassette mutagenesis, as shown below.

TX-108C-110C-2

101 102 103 104 105 106 107 108 109 110 111 112 113 114 115
G A T K L G E C T C D G S V Y
5'GC GCC ACA AAA TTA GGC GAA TGC ACT TGT GAT GGA TCC GTA TAT
3'-G TGT TTT AAT CCG CTT ACG TGA ACA CTA CCT AGG CAT ATA
Kasi | TX-108C-110C

TX-103b

116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131
D I Y R T Q R V N Q P S I I G T
GAT ATC TAC CGT ACC CAA CGC GTT AAT CAG CCA TCG ATC ATT GGA ACC
CTA TAG ATG GCA TGG GTT GCG CAA TTA GTC GGT AGC TAG TAA CCT TGG
XyTv-109

132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147
A T F Y Q Y W S V R R N H R S S
GCC ACC TTT TAT CAG TAC TGG AGT GTT AGA CGT AAT CAT CGG AGC TCC
CGG TGG AAA ATA GTC ATG ACC TCA CAA TCT GCA TTA GTA GCC TCG AGG
TX-108b

TX-154C-158C-162H-2

148 149 150 151 152 153 154 155 156 157 158 159 160 161 162 163
G S V N T A C H F N C W A Q H G
GGT TCG GTT AAT ACT GCA TGC CAC TTT AAT TGC TGG GCA CAG CAC GGG
CCA AGC CAA TTA TGA CGT ACC GTG AAA TTA ACG ACC CGT AGT GTG CCC
SphI TX-154C-158C-162H

164 165 166 167
L T L G
TTA ACC
AAT TGG GAT C
AvrII |

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F. Construction of TrX-DS8

[0062] The mutant TrX-DS8 was prepared using analogous methods as those outlined above in Sections A to E for the preparation of modified xylanases. TrX-DS8 incorporates the mutations found in N1-TX13 as disclosed in US 5,759,840. This mutations are N10H, Y27M and N29L. In addition, TrX-DS8 includes the following mutations: N44D, Q125A, I129E, Q162H and a disulfide bond between positions 110 and 154. The construction of the plasmid pTrX-DS8 was through ligation of overlapping phosphorylated oligonucleotides as described above.

[0063] Trx-DS8 exhibits the property of thermostability (Figure 4a), a pH profile paralleling that of TrX-162-DS1, and greater than 40% of optimum activity at physiological pH, and temperature.

Example 2: Characterization of mutant xylanases

A. Production of xylanases

[0064] The culture condition was identical to the well-established protocol described for other *E. coli*-expressed xylanases. A 5 ml of overnight inoculant in 2YT medium (16 g yeast extract, 10 g bacto-tryptone, 5 g NaCl, 1 L of water) containing ampicillin (100 mg/L) was added to 2YT medium (1 L) with ampicillin. The cultures were grown with shaking (200 rpm) at 37°C. After 16 hr, cells were harvested.

B. Purification of different disulfide bond-containing mutant xylanases

[0065] Protein samples were prepared from cells by first making an extract of the cells by grinding 10 g of the cell paste with 25 g of alumina powder. After grinding to smooth mixture, small amounts (5 mL) of ice cold buffer A (10mM sodium acetate, pH 5.5 for BcX mutants) or buffer B (10mM sodium acetate, pH 4.6 for TX mutants) were added and the mixture ground vigorously between additions. The alumina and cell debris were removed by centrifugation of the mixture at 8000 x g for 30 min.

[0066] The crude extract was heated at 60°C for 15 min and centrifugation to remove a large amount of precipitate. The supernatant was acidified to pH 4.6, frozen at -20°C overnight, thawed and centrifuged to remove more precipitate.

[0067] After the above pretreatment, the cell extract committed to column chromatography and was pumped onto a 50 mL bed volume, S-Sepharose fast flow, cation exchange column (Kabi-Pharmacia, Canada), equilibrated in buffer A. The xylanase was eluted with a 300 mL linear gradient of 0 to 0.3M NaCl in buffer A at a flow rate of 3 mL/min. The xylanase elutes at 100 to 150 mL of the gradient. The fractions are checked on SDS-PAGE, and those fractions having most of the xylanase were pooled, and concentrated by ultrafiltration using 3000 dalton molecular weight cutoff membranes (Amicon YM3). The concentrated material (5 mL) was then applied to a 1.5 cm x 85 cm TSK-HW50S gel filtration column, equilibrated in 50 mM ammonium acetate pH 6. The xylanase eluted at a volume of 90 to 100 mL. These fractions were analyzed by SDS-PAGE, and the peaks pooled as pure xylanase. The protein was quantified using the extinction co-efficient at 280 nm.

C. Standard assay for the measurement of enzymatic activity

[0068] The quantitative assay determined the number of reducing sugar ends generated from soluble xylan. The substrate for this assay was the fraction of birchwood xylan which dissolved in water from a 5% suspension of birchwood xylan (Sigma Chemical Co.). After removing the insoluble fraction, the supernatant was freeze dried and stored in a desiccator. The measurement of specific activity was performed as follows. Reaction mixtures containing 100 L of 30 mg/mL xylan previously diluted in assay buffer (50 mM sodium citrate, pH 5.5 or the pH optimum of the tested xylanase), 150 L assay buffer, 50 L of enzyme diluted in assay buffer were incubated at 40°C. At various time intervals 50 L portions were removed and the reaction stopped by diluting in 1 mL of 5mM NaOH. The amount of reducing sugars was determined with the hydroxybenzoic acid hydrazide reagent (HBAH) (Lever, 1972, Analytical Biochem 47:273-279). A unit of enzyme activity was defined as that amount generating 1 mol reducing sugar in 1 minute at 40°C.

[0069] For the comparison between mutant and the wild type xylanases (TABLE 3), the specific activities of a xylanase was converted to the relative activity which is its calculated in percentage as compared to the specific activity of the natural xylanase.

TABLE 3. Relative activity of TrX xylanases

Xylanase	Relative activity %
natl. TrX	100*

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(continued)

Xylanase	Relative activity %
TrX	103
TrX-DS1	116
TrX-162H-DS1	102
TrX-162H-DS4	91
* The specific activity of the natural TrX (770 U/mg) was normalized to 100%.	

[0070] As can be seen from Table 3, the specific enzymatic activities of the mutant xylanases at 40°C have not been changed significantly as compared to the natural xylanases.

Example 3: Thermostability of mutant xylanases

[0071] This was a test of the tolerance of xylanase to incubation at a set temperature, without any substrate. The xylanase (150 g/mL) in assay buffer (50 mM sodium citrate) was incubated at a set temperature or set period of time. Aliquots were cooled to room temperature (around 20°C), the residual enzymatic activity of all samples was determined via the HBAH assay at 40°C, as stated in Example 2C.

(A) Effect of length of incubation

[0072] The effect of the length of incubation on the activity of xylanase samples was determined at 62.5°C at pH 5.5 (Figure 3). Aliquots were removed at 0, 5, 10, 20, 30, 40 and 60 min for the determination of residual activity. The residual enzymatic activity at 0 min was normalized to 100%.

[0073] After 5 mins of incubation, the wild type TrX and the Q162H mutant TrX-162H (US 5,759,840) almost lost all residual activity, while the mutant TrX-DS1 with a disulfide bond, retained 60% of its residual activity. However, it retained only 20% of its activity at 20 mins and lost all activity at 40 min. In contrast, the mutant TrX-162H-DS1, with the additional mutation of Q162H, showed superior thermostability by retaining about 87% of its activity at 20 min, 78% at 40 min and 68% at 60 min. The mutant TrX-162H-DS4 with both 108/158 and 110/154 disulfide bonds retained 84% activity after 60 min.

(B) Effect of incubation temperatures on the residual activity of mutant TrX.

[0074] Thermostability of mutant TrX enzymes was also determined by tolerance of different incubation temperatures. Samples of xylanases were incubated in 50 mM sodium citrate buffer (pH 5.5) at different temperatures (48, 52, 56; 60, 64, 68, 70 and 72°C) for 30 min. The residual enzymatic activity of the samples was determined, with the residual activity at 48°C normalized to 100% (see Figures 4(a) and 4(b)). The T_{50} , which is the incubation temperature allowing the maintenance of 50% residual activity after 30 min, was determined for each mutant TrX.

[0075] Without wishing to be bound by theory, the higher T_{50} of TrX-162H-DS1 (65°C) versus TrX-DS1 (61 °C) demonstrates the enhancement of thermostability by the mutation Q162H in the disulfide mutants. The double disulfide mutant TrX-162H-DS4 also exhibited high stability with a T_{50} gain of +14°C over the natural TrX. Comparison of T_{50} of TrX-162H-DS1 (85°C) and TrX-162H-DS2 (59°C) indicates that the 110/154 disulfide in TrX-162H-DS1 contributes greater thermostability than the 108/158 disulfide in the latter. TrX-DS8 also exhibited high thermostability, with a T_{50} gain of +16 °C when compared to natural TrX.

(C) Effective incubation temperature

[0076] In the following example, a model study of the effect of the enzyme formulation on thermostability of the combination mutant was conducted in the presence of an additive, glycerol. The unmodified TrX and the mutant TrX xylanases were incubated for 30 min at 20, 50, 60, 70, 80 and 90°C in a buffer (pH 5.0) with 40% glycerol. The residual activity was determined by the HBAH assay. The residual enzymatic activity at 0 min was normalized to 100% (Figure 5).

[0077] At 50°C, all TrX samples retained their enzymatic activity. At 60°C, the wild type TrX retained 75% of its activity while TrX-DS1 and TrX-162H-DS1 retained 80 and 100% respectively (Figure 5). At 70°C, TrX-DS1 and TrX-162H-DS1 maintained 10 and 98% respectively. At 90 min, the latter retained 65% of the residual activity.

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(D) Effect of incubation time on the residual activity of TrX-162H-DS1 at 90°C

[0078] Sample of TrX-162H-DS1 in 40% glycerol and buffer were incubated in a covered circulating water bath (Haake type F 4391, with a fluctuation of 0.1°C) at 90°C. Temperature of the water bath was confirmed with a thermocouple. Aliquots were removed at 0, 5, 10 and 30 min for assay of residual activity. The residual enzymatic activity at 0 min was normalized to 100%.

[0079] At 5, 10 and 30 min, TrX-162H-DS1 retained 90, 85 and 65% of the residual activity respectively (Figure 6).

Example 4: Temperature/activity profile of mutant xylanases

[0080] This was a test on the effect on different temperatures to the enzymatic activity of the xylanase in the hydrolysis of soluble xylan. The procedure was identical to the standard assay (Example 2 C) with changes in the incubation temperature and time. The enzymes (1.5 µg/mL) and soluble xylanase in 50 mM sodium citrate buffer of pH 4.5 were mixed and incubated in a circulating water bath at different temperatures. After 30 min, the amount of reducing sugars released from xylan was determined by HBAH and was calculated as relative activity, with the value at temperature optimum as 100%.

[0081] The effect of temperature on the hydrolysis of xylan was shown in Figure 7. The natural TrX, TrX-DS1, TrX-162H-DS1, TrX-162H-DS2 and TrX-162H-DS4 enzymes all had the same temperature/activity profile, and the only difference is in the greater activity (80%) in mutant TrX-162H-DS4 as compared to the others (45%) at 60°C. These results indicate that the disulfide mutation, along with the Q162H mutation, has little or no effect on the optimal temperature (50°C) of TrX. In addition, all of the enzymes shown in the figure exhibit at least 40% of their optimum activity from about 40 to about 50 °C, which is suitable for feed pelleting applications.

Example 5: pH/activity profile of mutant xylanases

[0082] This was a test of the effect of different pH on the enzymatic activity of the xylanase in the hydrolysis of soluble xylan at the approximate physiological temperature of digesta.

[0083] The procedure was identical to the standard assay (Example 2 C) with changes in the incubation temperature and time. The *Trichoderma* enzymes natural TrX and mutant TrX (3.0 µg/mL) and soluble xylan in 50 mM sodium citrate buffers of pH 3-8 were incubated together at 40°C for 7 min. The amount of reducing sugars released from xylan was determined by HBAH and was calculated as relative activity, with the value at pH optimum as 100%.

[0084] The profile of the effect of pH on the enzymatic activity of TrX, TrX-162H-DS1 and TrX-162H-DS2 (Figure 8) are similar, thus indicating little or no effect of the mutations (disulfide bond formation and Q162H) on the pH optimum. The pH profile for TrX-DS8 was also similar to these modified xylanases (data not shown). All of the enzymes shown in the figure exhibit at least 40% of their optimum activity from about pH 3.5 to about pH 6, which is suitable for feed pelleting applications.

[0085] The double disulfide mutant TrX-162H-DS4 differed by showing slightly greater activity at the pH range higher than 6. At the acidic pH of 4-6 TrX, TrX-162H-DS1, TrX-162H-DS2 and TrX-162H-DS4 maintained at least 75% optimal activity.

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